Gastric antral vascular ectasia should not be overlooked in erythropoietin resistance: a series of case reports

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Background. Gastric antral vascular ectasia (GAVE) is currently recognized as an important cause of upper gastrointestinal (GI) haemorrhage, being responsible for about 4% of non-variceal upper GI haemorrhages and typically presents in middle-aged females. GAVE, also called "watermelon stomach", is diagnosed through esophagogastroduodenoscopy and is characterized by the presence of visible columns of red tortuous enlarged vessels along the longitudinal folds of the antrum. The pathogenesis is still obscure and many hypotheses have been proposed such as mechanical stress, humoral and autoimmune factors. In the last two decades, numerous therapeutic strategies have been proposed, including surgical, endoscopic, and medical choices, yet successful treatment of GAVE continues to be a challenge. Currently, given the rapid response, safety, and efficacy, endoscopic ablative modalities have largely usurped medical treatments as first-line therapy, particularly using argon plasma coagulation. The actual GAVE prevalence in patients with end-stage renal disease (ESRD) is not clear, yet in difficult cases it should be considered as a cause of erythropoietin resistance.

Case presentation. We report four clinical cases of GAVE syndrome patients diagnosed with stage 4 to 5 chronic kidney disease. All patients presented with anaemia and GI haemorrhage, the origin of which turned out to be GAVE syndrome.

Conclusions. GAVE syndrome is a serious condition in ESRD patients, especially in those presenting with treatment-refractory anaemia. Realization of its aetiology and characteristics is essential to suspect, diagnose, and treat gastric ectasia. Only proper diagnosis and well-timed disease treatment can significantly improve a patient’s medical condition and future prognosis.

Keywords: gastric antral vascular ectasia, GAVE, watermelon stomach, end-stage renal disease, ESRD, hemodialysis
INTRODUCTION

Gastric antral vascular ectasia (GAVE) syndrome, also known as “watermelon stomach”, is recognized as a rare cause of upper gastrointestinal bleeding (1) and can be a cause of severe blood effusion. Gastric antral vascular ectasia is often found with other disorders, such as liver cirrhosis, autoimmune connective tissue disorder, bone marrow transplantation, ischemic heart disease, valvular heart disease, hypertension, hypothyroidism, diabetes, or acute myeloid leukaemia (2–6). Additionally, recent studies show that patients with end stage renal disease (ESRD) are more likely to have GAVE syndrome (7). We report four clinical cases of GAVE syndrome in patients diagnosed with stage 2 to 5 chronic kidney disease undergoing continuous ambulatory haemodialysis. All patients presented with anaemia and GI haemorrhage, the origin of which turned out to be GAVE syndrome.

CASE REPORTS

Case 1
A 66-year-old female patient presented with complaints of general weakness and melena for five days. She had been on haemodialysis for three years after diagnosis of ESRD due to hypertension and diabetes. On examination, patient’s condition was stable, blood pressure (BP) 160/80 mmHg, heart rate 78 bpm. Rectal examination confirmed melena. As a result of continuous loss of blood in stool, laboratory investigations revealed iron-deficiency anaemia with haemoglobin value 60 g/l, despite continuous administration of intravenous iron and erythropoietin previously during each dialysis session. Upper gastrointestinal endoscopy showed multiple linear gastric vascular malformations with signs of bleeding. APC was performed to which the patient responded with positive dynamics of increased RBC (3.9 × 10¹²/l), haemoglobin (126 g/l), and platelets counts (268 × 10⁹/l). As the patient’s condition became stable, she was discharged on a conservative line of management.

Case 2
A 75-year-old male patient with hypertensive nephropathy on chronic haemodialysis for five years was referred to the hospital with complaints of general weakness and melena present for four days. His BP was 148/80 mmHg, heart rate 80 bpm. Laboratory investigations revealed slight iron-deficiency anaemia and haemoglobin value of 118 g/l. The patient also had a history of myocardial infarction, heart insufficiency and coronary artery bypass. Upper gastrointestinal endoscopy found multiple linear gastric vascular malformations in the antrum, with 3 mm lesions and no signs of bleeding. During the procedure,
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bleeding occurred and adrenaline injections were given. APC was performed for GAVE syndrome and the patient responded well, with positive dynamics showing increased red blood cell \((5.1 \times 10^{12}/l)\), haemoglobin \((136\text{ g/l})\) and platelets \((268 \times 10^9/l)\) counts.

Case 3

A 64-year-old male patient with ESRD due to chronic pyelonephritis, on maintenance haemodialysis for five years presented with complaints of general weakness and shortness of breath for four days in a row. On examination, the patient’s condition was unstable with weakened vesicular breathing sounds, peripheral leg oedema, severe pallor and indolence, BP 105/60 mmHg, heart rate 96 bpm. The patient also had a history of two myocardial infarctions, severe heart insufficiency, coronary artery bypass, and cardiac stimulator implantation surgery. Rectal examination was positive for melena and, as a result of continuous loss of blood in stool, laboratory investigations revealed iron-deficiency anaemia with haemoglobin of 98 g/l. As chest radiograph showed pleural effusion on both sides of the lungs, right lung puncture was performed with 1.5 l of transudate retrieved. After upper gastrointestinal endoscopy, the patient was found to have multiple linear gastric vascular malformations in the antrum without any signs of bleeding. Due to pleural effusion and peripheral leg oedema remaining during the hospital stay, the patient was given additional haemodialysis course. Despite maximal doses of erythropoietin therapy along with blood transfusion and intravenous iron in view of the severe iron deficiency, this patient remained anaemic. APC was performed, and one month after endoscopic therapy there was no need for transfusion of RBC, haemoglobin concentration stabilized. The patient responded well with increased blood

Fig. 3. Argon plasma coagulation treatment of GAVE

Fig. 4. “Watermelon stomach” – linear, intensely erythematous lesions at the apices of longitudinal antral folds radiating to the pylorus

Fig. 5. Small multiple linear gastric vascular malformations in the antrum with moderate bleeding
counts (red blood cell $3.6 \times 10^{12}$/l, haemoglobin 126 g/l) and was discharged on a conservative line of management.

**Case 4**
An 80-year-old male patient with CKD stage G4 of unknown origin was referred with complaints of general weakness, shortness of breath, and blood in stool for five days in a row, with severe worsening of the symptoms. He denied previous history of liver or autoimmune disease, and this was the first episode of melena. On physical examination, the patient’s condition was stable with notable mucosal pallor, shortness of breath, rhythmic heart activity, vesicular breathing sounds, and lack of peripheral oedema, BP 85/58 mmHg, heart rate 82 bpm. In the past, myocardial infarction and atrial fibrillation were diagnosed; heart ablation surgery and cardiac stimulator implantation were performed. Rectal examination was positive for melena and laboratory investigations revealed iron-deficiency anaemia with haemoglobin of 77 g/l. Urea blood level was 8.2 mmol/l, creatinine 190 µmol/l and glomerular filtration rate 28 ml/min/1.73 m². During hospital stay, the patient was treated with erythropoietin therapy, blood transfusion, and intravenous iron therapy. Endoscopy found multiple linear gastric vascular malformations in the antrum with small signs of bleeding. Argon plasma coagulation was performed for GAVE syndrome, to which the patient responded well with increased red blood cell ($4.8 \times 10^{12}$/l) and haemoglobin (123 g/l) counts. The altered blood in stool decreased and the patient was discharged on a conservative line of management.

<table>
<thead>
<tr>
<th>Table. Overview of presented case reports</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>CASE 1</strong></td>
</tr>
<tr>
<td>CKD grade</td>
</tr>
<tr>
<td>Haemodialysis</td>
</tr>
<tr>
<td>Complaints</td>
</tr>
<tr>
<td>Erythropoetin dose</td>
</tr>
<tr>
<td>Intravenous iron administration</td>
</tr>
<tr>
<td>Haemoglobin (before/after treatment)</td>
</tr>
<tr>
<td>Underlying conditions</td>
</tr>
<tr>
<td>Treatment</td>
</tr>
</tbody>
</table>
DISCUSSION

Although GAVE syndrome is involved in only 4% of non-variceal upper gastrointestinal bleeding developments (1), more and more case reports describe ESRD coexistence with GAVE syndrome (11). The way to suspect and anticipate the occurrence of this syndrome is very important, especially for patients with ESRD. If a patient has iron or erythropoietin resistance anaemia, undetected bleeding loss can quickly reduce the patient’s iron reserves (15). There are theories trying to explain the origin of the syndrome.

It has been suggested that in patients with chronic kidney disease, increased mechanical stress of the antrum in uraemia-induced weakening of gastric emptiness is thought to contribute to the development of GAVE. Enhanced gastrin levels also increase gastric vasodilatation (9). Uraemia-induced platelet dysfunction might also be incriminated as a risk factor for antral bleeding (10). Prostaglandin E2 is another hormone with vasoactive functions; in a single study, it was found to be significantly higher in cirrhosis patients with GAVE than in those without. In another study, histological examination showed that neuroendocrine cells containing 5-hydroxytryptamine and vasoactive intestinal peptide were found close to the ectatic gastric mucosal vessels (9). An impaired renal excretory function and decreased removal of these vasoactive substances from the circulatory system in patients with CKD may be associated with the emergence of gastric vascular ectasia (11–17). These pathological mechanisms were also noticed in the cases reported in the present paper. Three of the four patients suffered from stage 5D CKD and had been treated with haemodialysis three times per week for the past three to five years. The average time from the beginning of dialysis to the first bleeding episode was 52 months, although a larger sample size would be needed to be able to generalize this finding to the general population. However, every case should be approached individually as the time GAVE takes to manifest varies.

Typical initial presentations of the syndrome range from occult bleeding, causing transfusion-dependent chronic iron-deficiency anaemia, to severe acute upper gastrointestinal bleeding (8). All four described patients presented with anaemia levels between 60 and 118 g/l, as well as symptoms such as weakness, pallor, and shortness of breath. Three patients developed evident GI haemorrhage confirmed by rectal examination.

Literature suggests that lesions are typically located in the gastric antrum, and the four described cases were in line with this. Gastric lesions slightly raised above the mucosal surface, cherry red in colour and 2–10 mm in size were seen. GAVE syndrome appears as endoscopic forms of “watermelon stomach”, characterized by longitudinal flat rows and reddish stripes diverging from the pylorus into the antrum. Endoscopy findings were sufficient to diagnose GAVE syndrome and histological investigation was not necessary.

The initial form of treatment was APC (17, 18), which is considered the preferred and the most frequently used method of endoscopic therapy. In our four patients endoscopic haemostatic therapy using APC was performed for the ectatic lesions; the procedure was well tolerated without any further complications associated with the therapy. After the endoscopic procedure, GAVE lesions were significantly diminished. All patients required additional erythropoietin therapy, blood transfusions, and intravenous iron. After the therapy, patients’ condition became stable and they were discharged on a conservative line of management.

According to the documented cases of patients with GAVE syndrome undergoing APC therapy, appropriate monitoring is required as there is a risk of eventual recurrence. Thus, in recurrent bleeding, repeat endoscopy is useful and surgery is necessary.

CONCLUSIONS

In conclusion, GAVE syndrome is an important cause of upper gastrointestinal bleeding and has a higher prevalence in patients with chronic kidney disease. Verification of the aetiology and characteristics are essential for suspecting, diagnosis, and treatment of gastric ectasia in patients, especially in those with more than one risk factor and showing signs of treatment refractory anaemia. Only proper diagnosis and well-timed treatment can significantly improve patient’s medical condition and future prognosis.

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References

**Santrauka**

**Ivadas.** Skrandžio urvo kraujagyslių ektazija yra svarbi viršutinės virškinimo trakto hemoragijos priežastis, dėl kurios išsivysto maždaug 4 % viršutinio virškinimo trakto kraujavimų be varikozės. Paprastai tai pasireiškia vidutinio amžiaus moterims. Skrandžio urvo kraujagyslių ektazija, taip pat vadinama „Arbūzinu pilvu“, diagnozuojama ezofagogastroduodenoskopijos metu: endoskopiškai matoma ilgą kraščiagramą urve. Patogenezė vis dar yra neaiški, tačiau yra siūlomos hipotezės, tokios kaip mechaninis stresas, humoraliniai ar autoimuniniai veiksniai. Per paskutinius du dešimtmečius buvo pasiūlyta daugybė gydymo strategijų, tačiau sėkmingas skrandžio urvo kraujagyslių ektazijos gydymas vis dar yra iššūkis. Šiuo metu, atsižvelgiant į greitą reagavimą, saugumą ir veiksmingumą, endoskopinės abliacijos metodas yra pirmo pasirinkimo gydymas, ypač naudojant argono plazmos koaguliaciją. Skrandžio urvo kraujagyslių ektazijos paplitimas tarp pacientų, sergančių lėtine inkstų liga, ypač galutinės stadijos, néra aiškus, tačiau sudėtingose situacijose tai gali lemti atsparios gydymui eritropoetinu anemiją.

**Išvada.** Skrandžio urvo kraujagyslių ektazijos sindromas yra gyvybiškai pavojinga būklė, pacientams sergantiesiems toli pažengusia lėtine inkstų liga, ypač tiems, kuriem pasireiškia gydymui atspari anemija. Etiologijos ir ypatingų supratimas yra būtinas, norint įtarti, diagnozuoti ir gydyti skrandžio ektaziją. Tik tinka diagnozė ir savalaikis gydymas gali žymiai pagerinti paciento sveikatos būklę ir ateities perspektyvas.

**Raktažodžiai:** skrandžio urvo kraujagyslių ektazija, arbūzinis pilvas, galutinės stadijos lėtine inkstų liga, hemodializės

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**SKRANĘŽIO URVO KRAUJAGYSLIŲ EKTAZIJA – ATSPARIOS GYDYMUI ERITROPOETINU ANEMIJOS PRIEŽASTIS**

**Santrauka**

**Ivadas.** Skrandžio urvo kraujagyslių ektazija yra svarbi viršutinės virškinimo trakto hemoragijos priežastis, dėl kurios išsivysto maždaug 4 % viršutinio virškinimo trakto kraujavimų be varikozės. Paprastai tai pasireiškia vidutinio amžiaus moterims. Skrandžio urvo kraujagyslių ektazija, taip pat vadinama „Arbūzinu pilvu“, diagnozuojama ezofagogastroduodenoskopijos metu: endoskopiškai matoma išilginės kraujagyslių juostos prievarčio urve. Patogenezė vis dar yra neaiški, tačiau yra siūlomos hipotezės, tokios kaip mechaninis stresas, humoraliniai ar autoimuniniai veiksniai. Per paskutinius du dešimtmečius buvo pasiūlyta daugybė gydymo strategijų, tačiau sėkmingas skrandžio urvo kraujagyslių ektazijos gydymas vis dar yra iššūkis. Šiuo metu, atsižvelgiant į greitą reagavimą, saugumą ir veiksmingumą, endoskopinės abliacijos metodas yra pirmo pasirinkimo gydymas, ypač naudojant argono plazmos koaguliaciją. Skrandžio urvo kraujagyslių ektazijos paplitimas tarp pacientų, sergančių lėtine inkstų liga, ypač galutinės stadijos, néra aiškus, tačiau sudėtingose situacijose tai gali lemti atsparios gydymui eritropoetinu anemiją.

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